(;,	
	Please replace the paragraph beginning at page 19, line 17, with the following rewritten paragraph:
	-An immunodominant epitope of human thrombopoietin includes amino acids
	318 to 332 and has the following sequence (represented in single letter code):
a^3	LNTSYTHSQNLSEQ (SEQ ID NO:1)-
	Please replace the paragraph beginning at page 22, line 7, with the following rewritten
	paragraph: Analysis of the amino acid sequence of human thrombopoietin resulted in the
	identification of a predicted immunodominant epitope using the service provided by Epivax, Inc.
	Fourteen out of 15 amino acid residues of the immunodominant peptide identified using
4	antibodies from naïve patients showed 100% homology with the 14 residues at the C-terminal of
U	the 20 amino acid region predicted by EpiVax. The sequence of the predicted immunodominant
	epitope has an amino acid sequence identical to amino acid residues 312 to 331:
	TPTSPLLNTSYTHSQNLSQE (SEQ ID NO: 2)-
1	Please replace the paragraph beginning at page 40, line 8, with the following rewritten
	paragraph:
	A specific example of a modified polypeptide is a modified thrombopoietin
	having reduced immunogenicity while retaining substantial therapeutic activity. An
105	immunodominant epitope in native sequence human thrombopoietin is a c-terminal peptide
V	including amino acids 318 to 332:
	LNTSYTHSQNLSQEG (SEQ ID NO.: 1)-

Please replace the table beginning at page 50, line 14, with the following rewritten

paragraph;

-Table 2

Synthetic TPO c-terminal epitope peptides

and corresponding rabbit anti-peptide antibodies

1,6510

peptide sequence (length)	antibody ID#
154-170(17)	24
175-190(16)	48
195-211(17)	28
18-234(17)	19
244-259(16)	17
258-268(11)	49
268-283(16)	16
296-311(16)	51
318-332(15)	15

The peptides have the following sequences:

al

154- 170	RAPPTTAVPSRTSLVLT (SEQ ID NO: 3)
175-190	PNRTSGLLETNFTASA (SEQ ID NO: 4)
195-211	SGLLKWQQGFRAKIPGL (SEQ ID NO: 5)
218-234	SLDQIPGYLNRIHELLN (SEQ ID NO: 6)
244-259	SRRTLGAPDISSGTSD (SEQ ID NO: 7)
258-268	SDTGSLPPNLQ (SEQ ID NO: 8)
268-283	QPGYSPSPTHPPTGQY (SEQ ID NO: 9)
296-311	VVQLHPLLPDPSAPTP (SEQ ID NO: 10)
318-332	LNTSYTHSQNLSQEG (SEQ ID NO: 1)5

Please replace the paragraph beginning at page 52, line 12, with the following rewritten paragraph:

The entire linear amino acid sequence of human recombinant TPO was provided to EpiVax, Inc. for analysis and prediction of immunodominant epitopes. The results from EpiVax, Inc. identified 3 regions within the c-terminal epitope that have class II MHC binding motifs. One of these predicted epitope regions (identical to amino acids 312-331) is shown:

TPTSPLLNTSYTHSQNLSQE (SEQ ID NO: 2)-

Please replace the paragraph beginning at page 53, line 17, with the following rewritten paragraph:

L-A cDNA corresponding to the hTPO entire open reading frame was obtained by PCR using the oligonucleotide primes of the following Table.

CHO Expression

Vector PCR Primers

Cla.FL.F2 5' ATC GAT ATC GAT AGC CAG ACA CCC CGG CCA G 3'

(SEQ ID NO:11)

ORF.Sal 5' AGT CGA CGT CGA CGT CGG CAG TGT CTG AGA ACC 3'

(SEQ ID NO:12)

PRK5-hmpl I was used as template for the reaction in the presence of pfu DNA polymerase (Stratagene). Initial denaturation was for 7 min. at 94°C followed by 25 cycles of amplification (1 min. at 94°C, 1 min. at 55°C and 1 min. at 72°C). Final extension was for 15 min. at 72°C. The PCR product was purified and cloned between the restriction sites Clal and Sall of the plasmid pSV15.ID.LL to obtain the vector pSV15.ID.LL.MLORF.The sequence of the construct was verified.